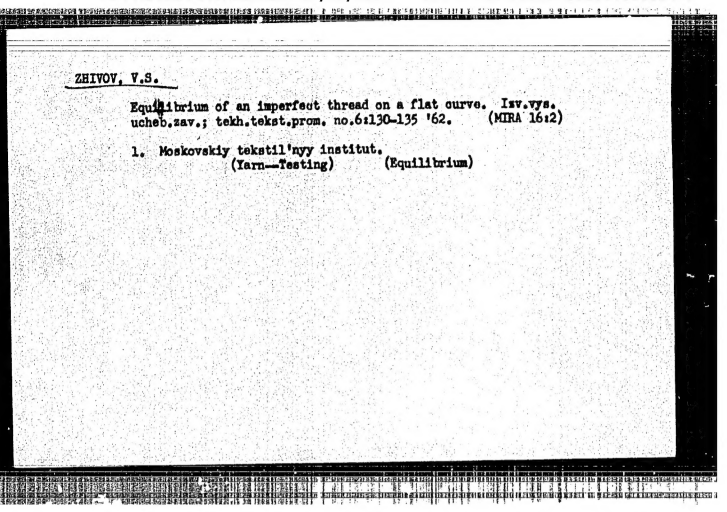
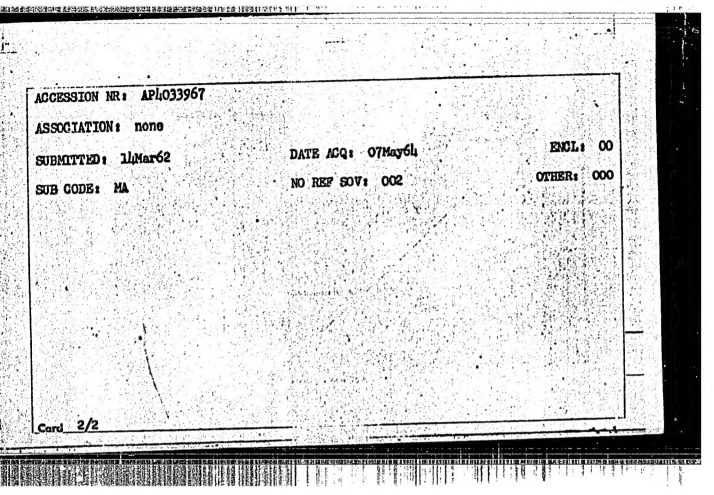


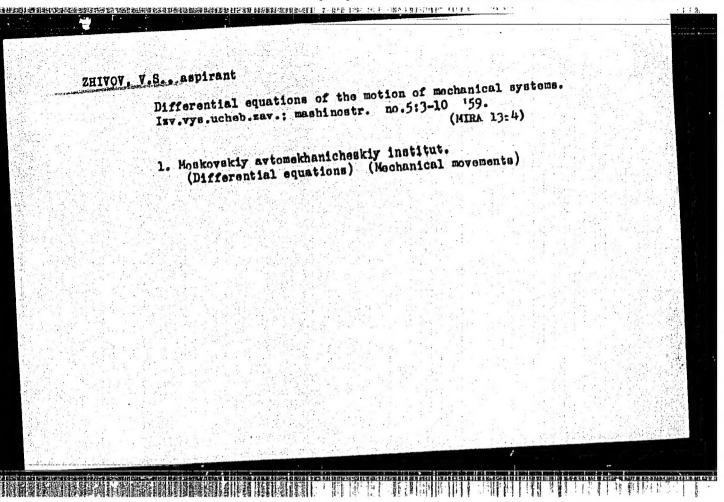
ZHIVOV, V. S., Cand Phys-Math Sci -- (diss) "Motion of combined systems." Moscow, 1960. 5 pp; (Ministry of Education RSFSR, Moscow Oblast Pedagogical Inst im N. K. Krupskaya); 200 copies; price not given; (KL, 26-60, 130)

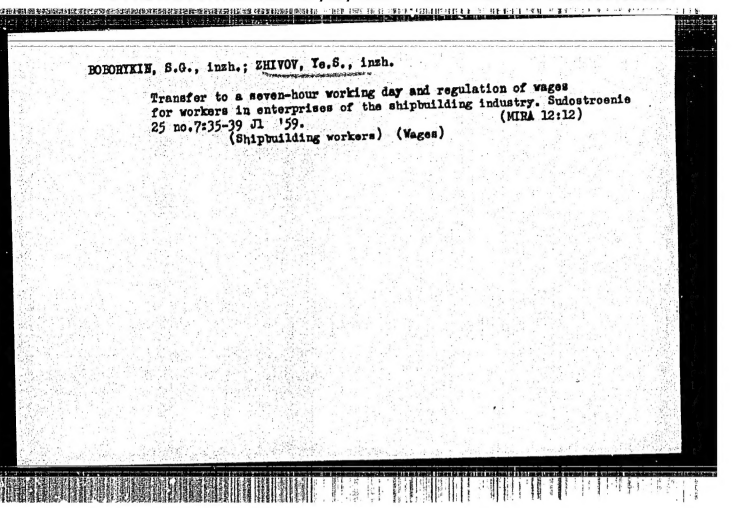


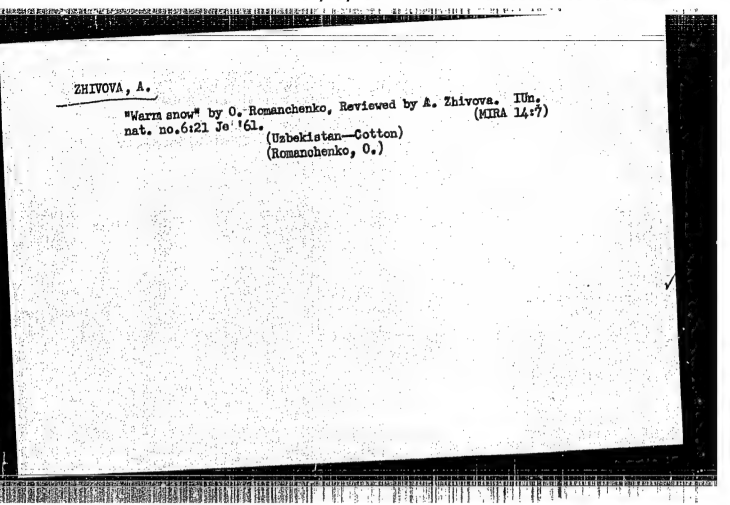
CCESSION NR: AP4033967	8/01110/611/000/002/0079/0081
WITHOR: Zhivov, V. S.	
ITLE: Stability of trajectorie	38
SOURCE: IVUZ. Matematika, no.	2, 1964, 79-81
ropic TAGS: stable trajectory, solution, perturbed integral cu	asymptotically stable trajectory, Lyapunov zero .
ABSTRACT: The author gives a r stable trajectories; intuitive closeness of a perturbed integr study of stability of trajector	easonable definition of stable and asymptotically ly stability of a trajectory means sufficient al curve to the unperturbed one. He reduces the iss of system
$\frac{dq_i}{dt} = Q_i (t, q)$	$(1, \dots, q_n), i=1,\dots, n$
to the study of stability of the equations, and gives an example	ne Lyapunov zero solution of a related system of o. Orig. art. has: 6 formulas.
Card 1/2	

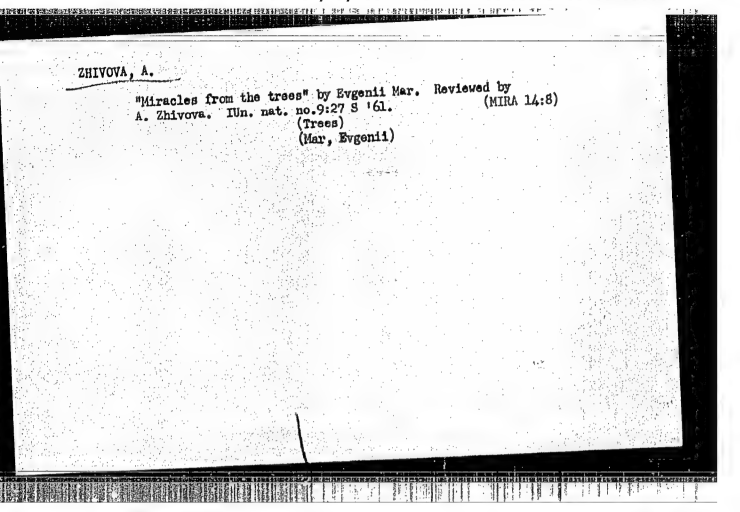
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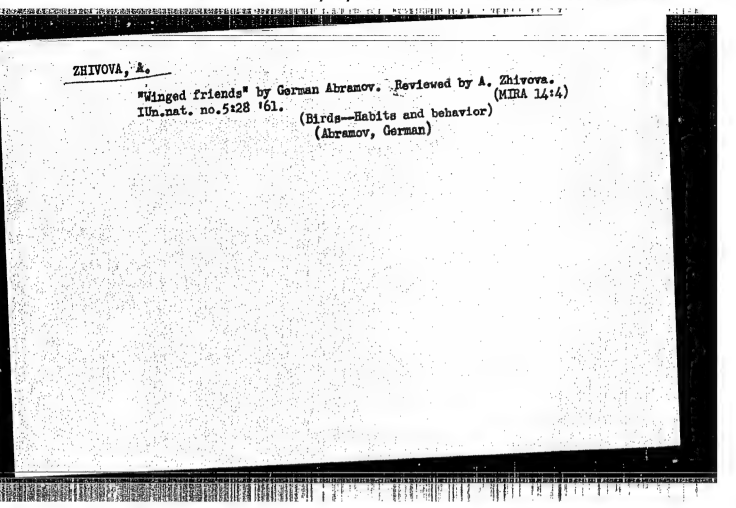


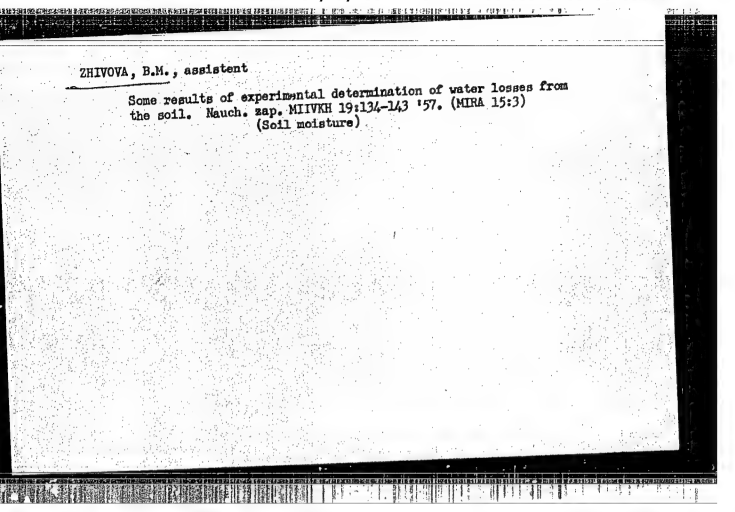


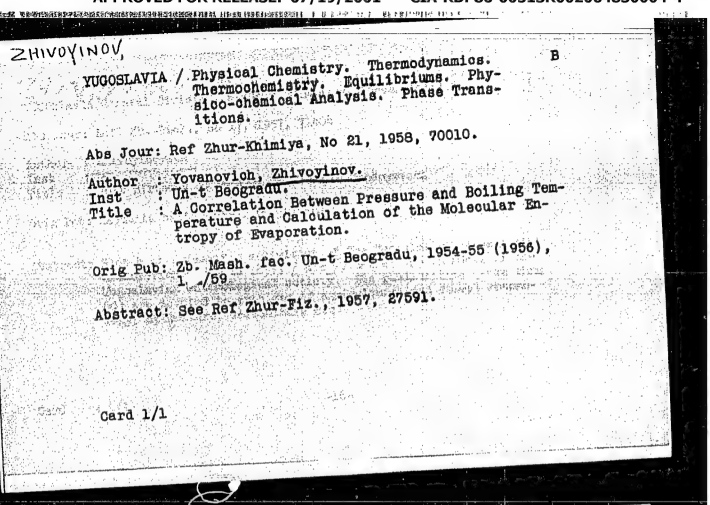


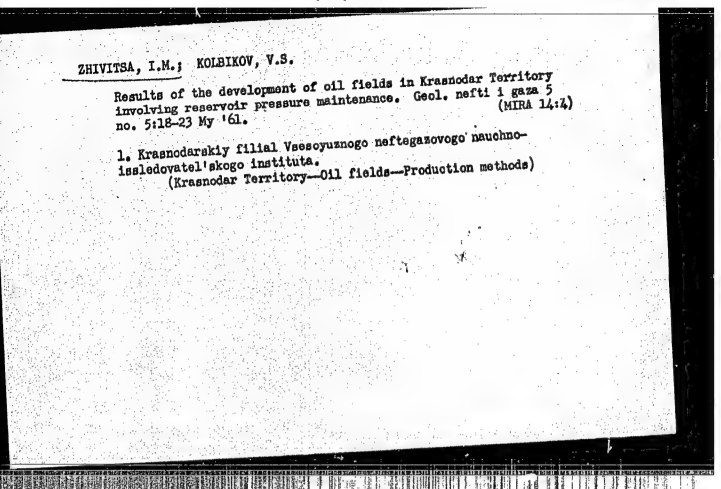


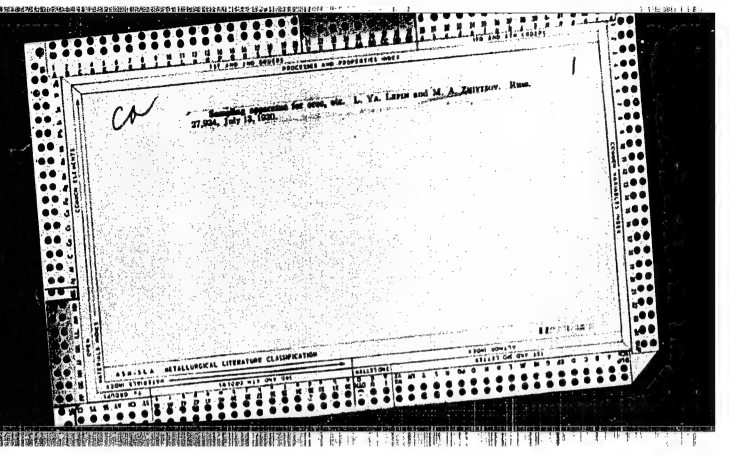


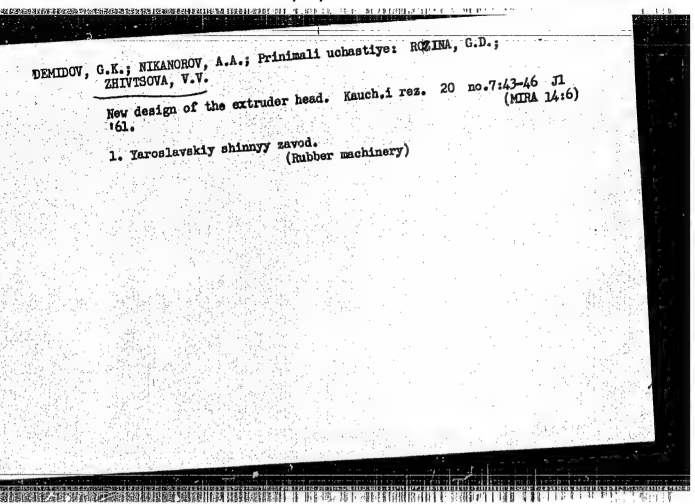




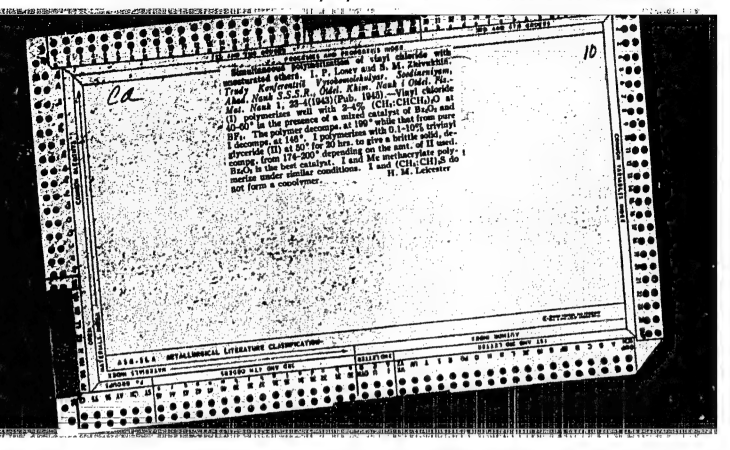






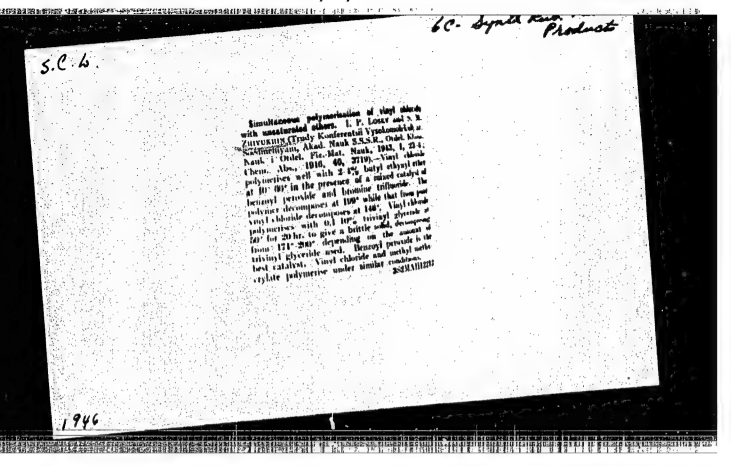


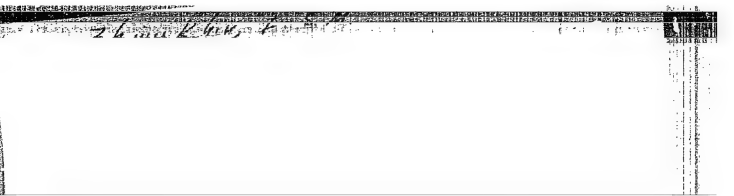
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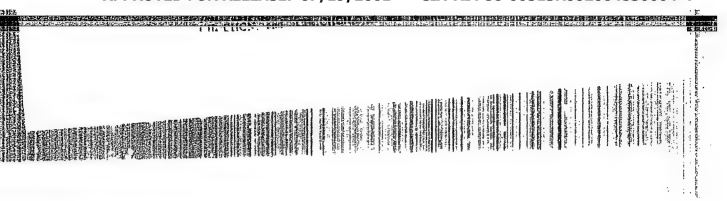
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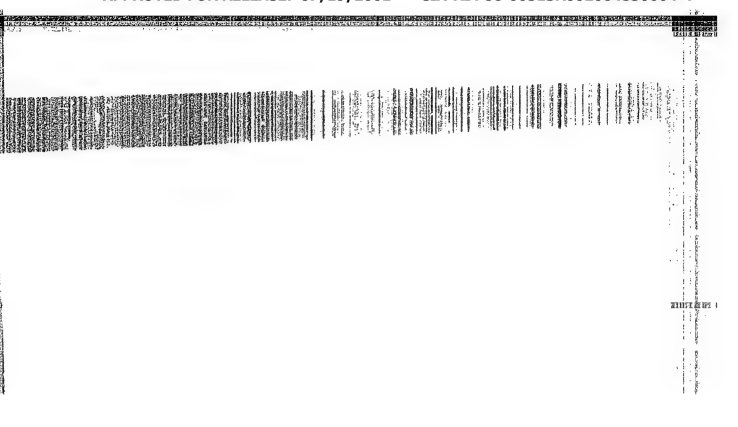
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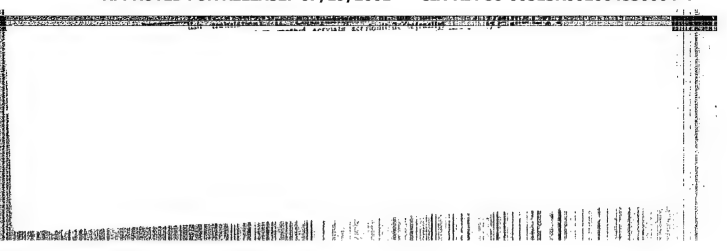


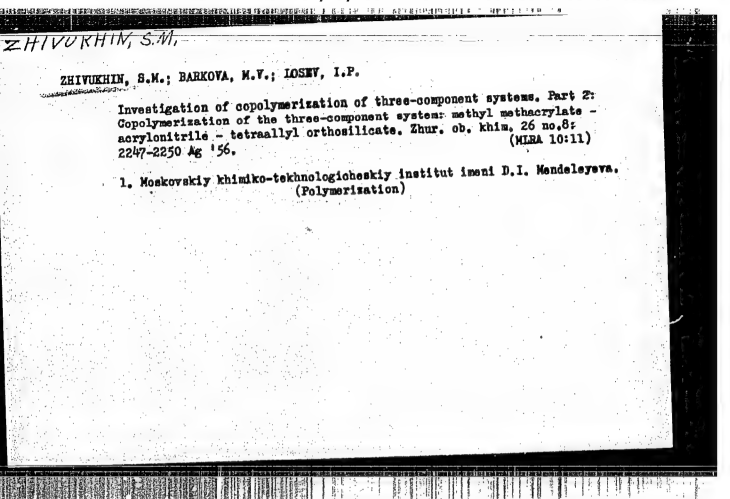












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S/191/60/000/008/005/014 B004/B056

AUTHORS:

PERIODICAL:

Zhivukhin, S. M., Dudikova, E. D.

TITLE:

Silanols and Their Transformations. Report I. Diphenylsilanediol and Its Transformation

Plasticheskiye massy, 1960, No. 8, pp. 23-25

TEXT: The authors first give a brief review of studies on the synthesis and transformations of silanols. They mention papers by N. N. Sokolov, V. A. Astakhin, M. M. Kotton, K. A. Andrianov, and B. N. Breytman, N. S. Nametkin, A. V. Topchiyev and F. F. Machus, M. F. Shostakovskiy, and D. A. Kochkin. The authors aimed at giving a precise description of the conditions for the synthesis of silanols, the study of their transformations and their combination with other organic substances. Diphenyltions and their combination with other organic substances. Diphenyltions and their combination with other organic substances. Diphenyltions of diphenyldichlorosilane in a mixture of toluene, water and isolysis of diphenyldichlorosilane in a mixture of toluene, water and isolysis of diphenyldichlorosilane in a mixture of toluene, water and isolysis of diphenyldichlorosilane in a mixture of toluene, water and isolysis of the white diphenylsilanedicle crystals fluctuates between 132 point of the white diphenylsilanedicle crystals fluctuates between 132 and 148 C because of inseparable anhydride forms, and increases to 152 C

Card 1/2

Silanols and Their Transformations. Report I. S/191/60/000/008/005/014 Diphenylsilanediol and Its Transformation B004/B056

because of condensation when stored for a length of time. The condensation of diphenylsilanediol was carried out at 150°C (molecular weight, 780-4080), 190°C (molecular weight, 2200-4650), and 230°C (molecular weight, 2870-4890). The melting points were between 30° and 60°C. Further, this compound was combined with: 1) with adipic acid (5 h at 190°C; ratio becompound was combined with: 1) with adipic acid (5 h at 190°C; ratio becompound was combined with: 1). The acid number decreased from 331 to tween the components, 1: 1). The acid number decreased from 331 to 147.5, and a brittle yellowish substance with a molecular weight of 1470. Was obtained. 2) With azelaic acid (1: 1, at 190° and 230°C), a 1390 was obtained. 2) With azelaic acid (1: 1, at 190° and 230°C), a yellow wax-like substance formed which had a molecular weight of 1470. Yellow wax-like substance was 3) With sebacic acid at 190° and 230°C, a brown wax-like substance was formed. With succinic and glutaric acids no stable products were obtained. The solubility differing from the components and the analyses confirmed the formation of condensates, and not of mixtures of the individual components. There are 16 references: 6 Soviet, 7 US, and 3 German.

Card 2/2

87646

s/191/60/000/012/005/016 B020/B066

15,9209

2209

Zhivukhin, S. M., Tolstoguzov, V. B.

AUTHORS: TITLE:

Phosphonitryl Chloride, Its Synthesis, Properties, and Use. Report No.1. Synthesis of Phosphonitryl Chloride

PERIODICAL: Plasticheskiye massy, 1960, No. 12, pp. 14 - 16

This is a survey of publications dealing with phosphonitryl chloride (PNC). It polymerizes to give a rubber-like material - a socalled "inorganic rubber". The course of this polymerization, the properties and the application of this material are described in recent publications and patents. PNC was later on obtained from PC15 and NH4C1, but in poor yields. Yields can be considerably increased when performing the reaction in an autoclave, or using tetrachloro ethane as solvent, or applying a protective layer of NH<sub>4</sub>Cl. In some variants of this method, a 52.6% yield of the trimer and a 25% yield of the tetramer, referred to theoretical yields, could be obtained. By means of the two newest methods suggested in 1957 also high yields are obtained. One of them is based

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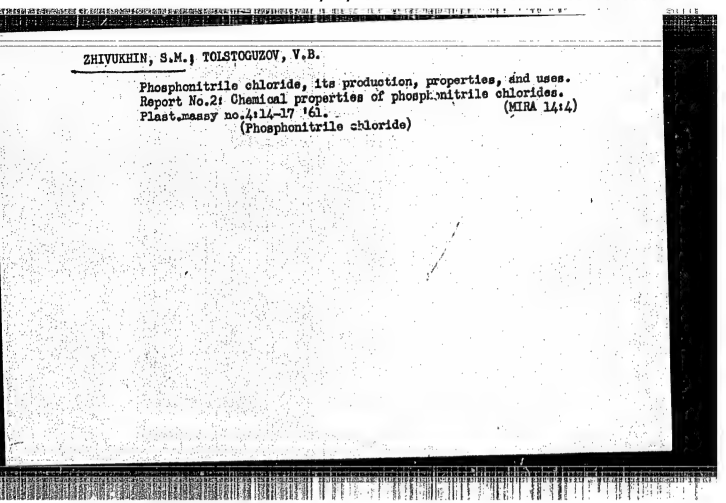
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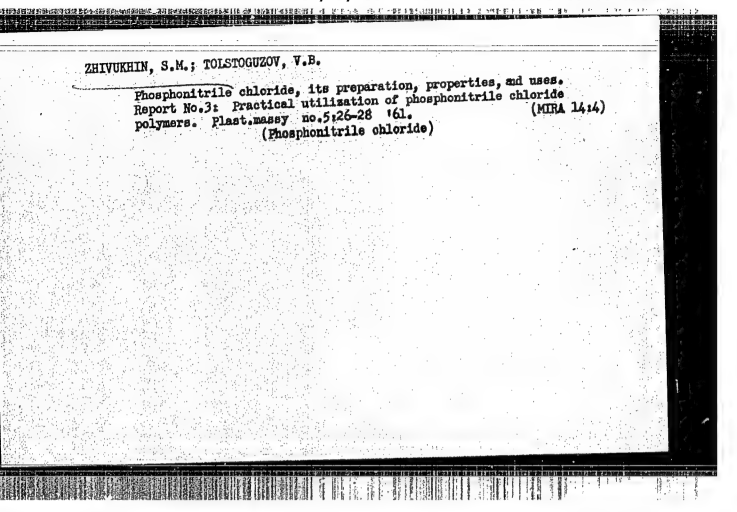
Phosphonitryl Chloride, Its Synthesis, Properties, and Use. Report No.1. Synthesis of Phosphonitryl Chloride

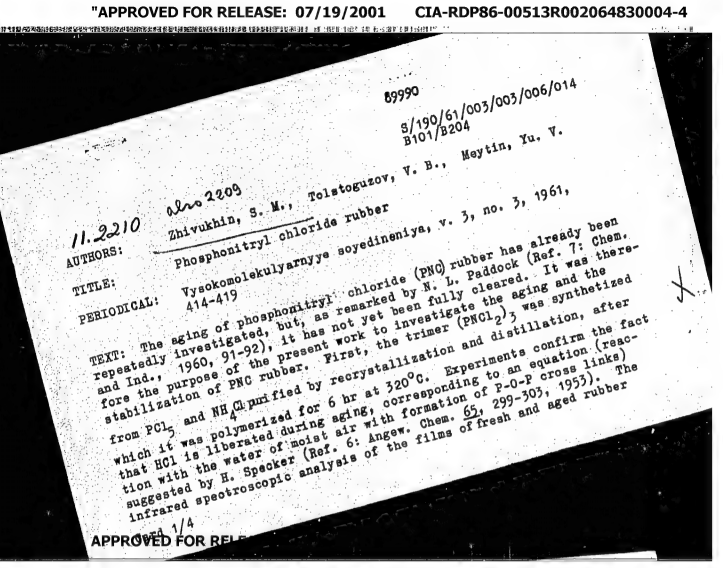
S/191/60/000/012/005/016 B020/B066

upon the reaction of NH4Cl with PCl5 in tetrachloro ethane in the presence of quinoline as catalyst, and yielded 35 - 40% trimer, 55 - 60% heptamer, and 0 - 5% tetramer and other polymers. The other method bases upon the reaction of a solution of PCl<sub>5</sub> in methylene chloride with liquid ammonia, in which a mixture of the trimer and tetramer and 0 - 5% of other polymers were obtained. The physiological properties of the product are given, the principal features of the separation of the polymer homologs and their purification are briefly described. The analytical methods for the resultant products are briefly described, and the results obtained by the laboratory of the kafedra organicheskikh i elementoorganicheskikh vysokomolekulyarnykh soyedineniy, MKhTI imeni Mendeleyeva (Department of Organic and Elemental-organic High-molecular Compounds of the Moscow Institute of Chemical Technology imeni Mendeleyev) with respect to PNC synthesis and the properties of the resultant reaction products are given. There are 47 references: 1 Soviet, 9 US, 12 German, 23 French, 3 British, and 1 Japanese.

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89990

Phosphonitryl chloride rubber

s/190/61/003/003/006/014 B101/B204

applied to KBr by means of a UR-10 spectrograph confirms; 1) The occurrence of a band corresponding to the P-O-P bond during aging. 2) The decrease of the intensity of the P=N band and occurrence of NH bands as a result of destruction. Furthermore, a displacement of the P=N bands with increase ing molecular weight from 1340 cm-1 (molecular weight 2.103) to 1360 cm (molecular weight 1.106) was observed. Pycnometrically, an increase in specific weight (from 1.77 to 2.02) was found, which takes place within 8 days. By X-ray analysis, this effect could be explained as crystallization. Conforming with the data of the increase in specific weight, a duration of the crystallization of 170 hr was found. The melting point of the crystals was between 30 - 40°C. By weighing, the effect produced by aging upon weight was investigated. With 100% moisture, the increase in weight was 1.4% after 200 hr, and 7.9% after 300 hr. This change in weight, however, depended on the ratio between the cross section of the specimen and its surface. The authors found: y = (P - P)/P = -0.111F - 0.7948 + 1.22 (4). Here, P is the initial weight, P after Thr, F the area of the cross section, cm2, & the thickness in mm. From this equation it follows that aging is a diffusion process proceeding from Card 2/4

89990

Phosphonitryl chloride rubber

S/190/61/003/003/006/014 B101/B204

the surface. 40 stabilizers were examined and the degree of stabilization K was determined:  $K = y_p/(y_p - y_{stab})$ , where  $y_p$  is the loss in weight of pure rubber according to Eq. (4), and  $y_{stab}$  is the loss in weight of stabilized rubber. Table 2 shows the results obtained by means of some stabilizers. There are 5 figures, 2 tables, and 11 references: 1 Soviet-bloc and 10 non-Soviet-bloc. The 1 reference to English-language publication is given in the text of the abstract.

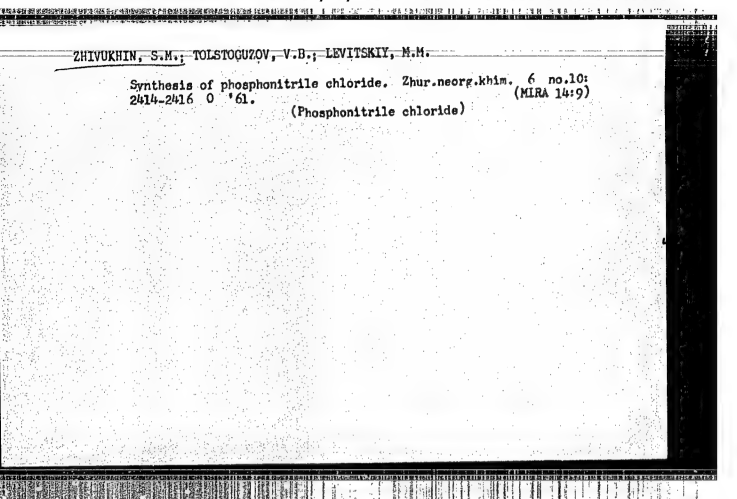
ASSOCIATION: Khimiko-tekhnologicheskiy institut im. D. I. Mendeleyeva

(Chemotechnical Institute imeni D. I. Mendeleyev)

SUBMITTED: July 7, 1960

Card 3/4

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				89990	4 /003 /0	02/006/014	
uo a buc	onitryl chloride rub	10 <b>6</b> F		B101/B2	204	03/006/014	X
	Наименование стабилира-	Количество		Наименование стабилива-	Количество		
	тора или циполнители	стабиднаа- Суора, %		TODA MAN MAGOVMOTORS	стабилиза. Огора, %	K	
	<b>Э</b> Спликоновый научун	4,1	0,98	ПолинаргомикоП	6,8	0.89	
	У Стеарат бария О Стеарат олова	3,7	0,93	Sb <sub>2</sub> O <sub>8</sub>	5,0	0,90	
	<b>О</b> Ортоборная кислота			Cr₂O₃ CuO	43,0 40,0	0,89 0,81	
	() Стеарат кодиня () Поливиниловый синрт	2,8	0,89	PbO <sub>s</sub>	30,0	0,72	
	Broumminionin cunbt	8,0 (	0,80	Sb <sub>1</sub> O <sub>8</sub>	43,0	0,76	
أأبيان برم إحيستنج						-4-14-14	
	to Table 2: 1) Sta	cium stes	rate	iller. 2) Quant	te. 6)	Ortho-bori	0
egend	con rubber. A) Bar	, _ ~ ~ , ~ ~ ~ ~ ~	alvy	invi alcohol.	) Polyu	rea.	
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) Sili	lcon rubber. 4) Bai	te. 8) P	01,7 4				



27511 s/079/61/031/009/012/012 D215/D306

AUTHORS:

Zhivukhin, S.M., Dudikova, E.D., and Kireyev, V.V.

TITLE:

Synthesis and study of organostannoxanes. I.

PERIODICAL: Zhurnal obshchey khimii, v. 31, no. 9, 1961, 3106 - 3111

TEXT: The present work was conducted to investigate the possibility of producing polyorganostannoxanes from certain dialkyltin dichlorides of the general formula R<sub>2</sub>SnCl<sub>2</sub> where R = C<sub>2</sub>H<sub>5</sub> and C<sub>3</sub>H<sub>7</sub>. The starting materials were obtained by K.A. Kocheshkov's method (Ref. 9: Sinteticheskiye metody v oblasti metalloorganicheskikh soyedineniy. Izd. AN SSSR, v. 5, 1947). Alkyltin acetates were prepared by reacting the corresponding dialkyltin dichloride with an acetate in an aqueous medium. Under such conditions, however, the acetates formed decomposed giving products of hydrolysis and condensation, i.e. compounds of the type

Card 1/7

27511 S/U79/61/031/009/U12/012 D215/D306

Synthesis and study of ...

[**周周武治·郑** 张宏宏说话在我的基础主动证式。它是没有被称为自然并指示于欧斯特。他们是自由自己的证明,但是自由自己的证明。

CH,COO  $\begin{bmatrix} R & R \\ SnO & SnOCOCH_{ii} \end{bmatrix}$ , where n = 2, 3

The reaction was carried out at 0°, 20° and 50°C. Compounds obtained at 50°C did not dissolve completely in benzene due to the admixtures of diethyltin oxide, but the yield corresponded to maximum. Compounds obtained at 0 and 20°C were soluble in organic solvents. To prevent the formation of diethyltin oxide the reaction was carried out at pH < 7. Preparation of polypropylstannoxanes is most suitable conducted at 50°C when up to 70% yields are obtained. Further experiments were concerned with increasing the molecular weight of the acetates prepared by subjecting them to the action of moisture containing air, in vacuum at different temperature over a period of 3 - 32 hrs. The resulting products were waxy

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Synthesis and study of ...

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solids soluble or partially soluble in organic solvents, some of their properties, in relation to the conditions of treatment, are given in Table 2. The molecular weight increases were, however, small and the treatment resulted mainly in decreases of solubility. The analysis of insoluble residue, from the treatment of polyethylstannoxane at 180-185°C, showed the increase of the oxygen content with simultaneous reduction of carbon and hydrogen contents. It may, therefore, be assumed that ethyl radicals bonding molecular chains were substituted by oxygen atoms, in the process involving the reaction of organic radicals linked to tin atoms with weak organic acids and the formation of the corresponding hydrocarbons. Acetic acid liberated during the condensation promotes separation of alkyl radicals and formation of acetate groups in the side chains of the polymeric molecules, followed by cross-linking and production of insoluble or slightly soluble compounds. Heating of the original alkylstannoxanes and their acetates in aqueous alkali brings about deeper transformations then the expected saponification of terminal acetate groups and the resulting products are in-

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Synthesis and study of ...

soluble in organic solvents. Prolonged heating in the presence of excess water resulted in the separation of the acetic acid

$$\begin{array}{c|c} CH_3COO & R & R \\ Sn-O & -Sn-OCOCH_3 + 2H_3O \rightleftharpoons \\ R & R \end{array}$$

$$\begin{array}{c|c} R & R \\ Sn-O & -Sn-OH + 2CH_3COOH \end{array}$$

The saponification products, dihydroxyalkylstannoxanes were white brittle solids with softening point of 200°C. They dissolved, with difficulty, in hot benzene and dioxane and the molecular weight of the soluble part corresponded to 900-1000. The presence of terminal hydroxyls opens the possibility of condensing these compounds with corresponding di- and poly-functional substances, e.g. polyalkylstannoxane-acetates. Polyethylstannoxane-acetates were pre-

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Synthesis and study of ..

pared by mixing aqueous solutions of diethyltin dichloride, sodium acetate and acetic acid at 20°C. After standing for 1.5 hours, the solid was filtered, washed and dried, the yield was 54-55 %, molecular weight 700-720. Polypropyl stannoxane was similarly prepared but at 50°C, by adding dipropyltin dichloride solution in methanol to aqueous acetate. The yield was 70.5 %, molecular weight 883-914. Dihydroxypolyethylstannoxane was obtained by refluxing polyethylstannoxane acetate with water for 5 hours. After neutralizing the acid the residue was boiled for 5 hours with more water and the procedure repeated several times. The yield of product was 60 %. The dihydroxypolystannoxane obtained was then condensed with an equimolecular quantity of polyethylstannoxane acetate at 160°C for 5 hours and 200°C for 5 hours. The product was a brown transparent mass slightly softening at 250°C. There are 3 tables, and 12 references: 6 Soviet-bloc and 6 non-Soviet-bloc. The four most recent references to the English-language publications read as follows: J.C. Montermoso, T.M. Andrews, L.P. Marinelli, J. of Polymer Sci., 32, 523, 1958; T.M. Andrews, F.A. Bower, B.R. LaLiberte, J.

Card 5/7

27511 S/079/61/031/009/012/012 D215/D306

Synthesis and study of ..

C. Montermoso, J. Am. Chem. Soc., 80, 4102, 1958; British Patent 718,393, 1954; R. Sasin, J. Org. Ch., 20, 770, 1955.

ASSOCIATION: Moskovskiy khimiko-tekhnologicheskiy institut im. D.I. Mendeleyeva (Moscow Institute of Chemistry and Technology im. D.I. Mendeleyev)

SUBMITTED: October 13, 1960

Table 2.

Legend: 1 - Starting compounds; 2 - reaction temperature; 3 - time of reaction (hrs.); 4 - solubility in benzene; 5 - polyethylstannoxane acetate (mol.wt. 796); 6 - 7 - polyethylstannoxane acetate (mol.wt. 796); 8 - polypropylstannoxane acetate (mol.wt. 898; 9 - polypropylstannoxane acetate (mol.wt. 898); 10 - total; 11 - cloudy solution; 12 - strongly cloudy solution, precipitate, precipitate; 13 - total; 14 - cloudy solution.

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33439 s/064/62/000/001/002/008 B110/B138

15.8180

Zhivukhin, S. M., Tolstoguzov, V. B. AUTHORS:

TITLE:

Production of phosphorus nitryl chloride

Khimicheskaya promyshlennost', no. 1, 1962, 19 -PERIODICAL:

TEXT: A method is developed for the industrial production of phosphorus nitryl chloride according to  $nPCl_5 + nNH_4Cl \rightarrow (PNCl_2)_n + 4nHCl + 120$ kcal/mole. The trimer (II) produced in the experiments contained tetramer (I) impurities, was crystalline (m. 108 - 114°C), had a slight odor and almost imperceptible irritant properties. Vapor pressures between 75.2 - 114.9°C and 114.9 - 189.3°C were determined from log P = 11.187 - 3979/T and log P = 8.357 - 2880/T, respectively. Heat of vaporization was 13.2 kcal/mole, heat of sublimation 18.2 kcal/mole, and heat of fusion 5.0 kcal/mole. The eutectic mixture II (0.65 - 0.7 molar parts) + I melts at 89.0 - 89.5 C. Best solubility of II is in benzene (55.0 g/100 g), and of I in CCl<sub>4</sub>. Molecular weight of II is 340 - 450 8. It decomposes in

moisture, is not corrosive, and can be stored for ever. The resulting

APPROVED FOR RELEASE: 07/19/2001

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Production of phosphorus nitryl...

dark yellow-to-brown oily liquid ( $D_{20} = 2.01$ ; at  $60^{\circ}$ C,  $\eta = 110$  cp; n<sub>D</sub><sup>14.5</sup> = 1.5857; MW = 1172 - 1392; 24.9% P; 59.63% Cl; 11.74% N; mean degree of polymerization: 10 - 12) is a mixture of higher polymer homologs  $(PNCl_2)_n$ , n > 4, soluble in benzene, toluene, xylene, acetone, and insoluble in water, acetic acid, and petroleum ether. The higher homologs have better solubility in benzene than the lower ones. The optimum solvent chlorobenzene (1.65 - 1.75 liters/kg of PCl5) melts near the optimum boiling point (128 - 130°C) causing violent agitation. The optimum molar ratio of NH<sub>A</sub>Cl to PCl<sub>5</sub> is 1.15 - 1.17. Optimum synthesis period is 8 - 12 hrs in the presence of 0.160 - 0.163 moles of quinoline per mole of PCl<sub>5</sub>. The trimer yield decreases and the mean polymerization degree increases in 12 hrs. The reaction is at first violent, bulk conversion takes place after 7 - 8 hrs, and then HCl is separated. The starting materials are put into enameled vessel 2 (Fig. 2) with reflux condenser 3. The separated HCl gas is collected in 5, the reaction mixture is cooled to room temperature, and quinoline hydrochloride and the  $\mathrm{NH}_A\mathrm{Cl}$  excess are

33439 5/064/62/000/001/002/008 B110/B138 Production of phosphorus nitryl.. filtered off at 6. C6H5Cl is distilled off in 8 and recycled. the oily residue is separated into phosphorus nitryl chloride and oil. The crystals are dissolved in petroleum ether, and the saturated solution is passed through filter 16. Partial distillation of the petroleum ether occurs at 18, and the trimer is crystallized in 20. It is centrifuged at 21, and the mother liquid passes into 18 again. The purified trimer (42% related to PCl5) is separated from the tetramer by fractional vacuum distillation, brystallization, or by CH3OH. The oil is passed to 22 (Fig. 3), dissolved in benzene, precipitated by petroleum ether, and left standing for 5 hrs. The bottom layer is pure oil, and the top layer is a mixture of solvents containing impurities. It is distilled at 80 - 100°C in 24. The bil yield is 40 - 42%. Quinoline hydrochloride and NHACl from 6 (Fig. 2) are rendered weakly alkaline in 27 (Fig. 4) by 25% NH,OH: For extraction, benzene is added by N·HC1 Card 3/0

33439 \$/064/62/000/001/002/008 B110/B138

Production of phosphorus nitryl....

stirring, and the mixture is left standing for 2 - 3 hrs. The solution of quinoline in benzene is passed to 29, dried with KOH for 24 hrs, filtered and distilled in 30. For 1 kg of phosphorus nitryl chloride, 1.96 kg of PCl<sub>5</sub>, 0.59 kg of NH<sub>4</sub>Cl, and 0.19 kg of quinoline are used.

There are 4 figures, 2 tables, and 5 references: 2 Soviet and 3 non-Soviet. The two references to English-language publications read as follows: H. N. Stokes, Am. Chem. J., 19, 782 (1897). N. L. Paddock, Brit. Plast., 31, no. 11, 473, 494 (1958).

Fig. 2. Flow chart for phosphorus nitryl chloride production (production and separation of polymer homologs).

Legend: (a) vapor; (b) vacuum; (c) brine; (d) nitrogen; (e) water; (f) HCl; (g) trimer for purification; (h) Fig. 3; (i) chlorobenzene; (k) quinoline; (l) petroleum ether; (m) oil.

Fig. 3. Purification of the oily liquid.

Legend: (a) of 12 and 16, Fig. 2; (b) petroleum ether; (c) benzene; (d) solvent for rectification; (e) vapor; (f) vacuum; (g) oil.

Card 4/0 4

S/078/62/007/009/004/007
B144/B101

AUTHORS: Zhivu!:hin, S. M., Tolatoguzov, V. B., Ivanov, A. I.

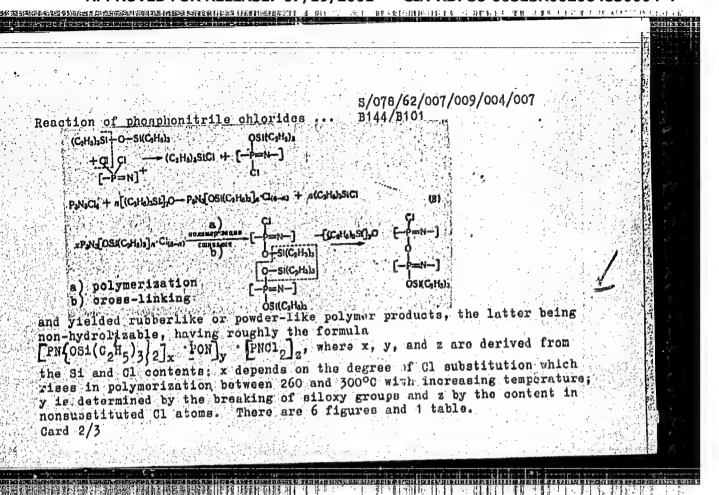
TITLE: Reaction of phosphonitrile chlorides with silanols, silanolates, and hexaalkyl disiloxane

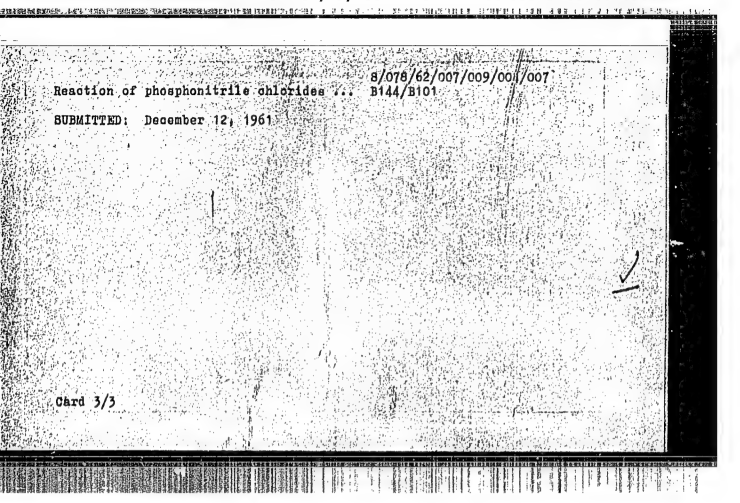
PERIODICAL: Zhurnal neorganicheskoy khimii, v. 7, no. 9, 1962,2192-2199

TEXT: Tests with triethyl and triphenyl silanoles and Na silanolates were unsuccessful but proved that PNCl<sub>2</sub> attacks the Si-0-Si bond. Hence hexaethyl disiloxane which contains one Si-0-Si bond was made to react at 230°C with PNCl<sub>2</sub> trimer (molar ratio 6:1). Substitution was obtained.

APPROVED FOR RELEASE: 07/19/2001 CIA-RDP86-00513R002064830004-4"

Card 1/3





5/079/62/032/009/011/011 1048/1242

AUTHORS:

Zhivukhin, S.M., Dudikova, E.D., and Ter-Sarkisyan, E.M.

TITLE:

Synthesis and investigation of organostannoxanes. II

PERIODICAL: Zhurnal obshchey khimli, v.32, no.9, 1962, 3059-3061

TEXT: The first paper on this subject appeared in this publication, v. 31, 1961, 3106. This paper reports the results of an attempt to prepare polyorganestrannoxanes by polycondensation of dibutyldiace-toxystannane and dibutyldibutoxystannane according to the equation

The experiments were carried out in an inert gas stream with constant stirring at 150 to 200°C. The starting materials were syn-

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APPROVED FOR RELEASE: 07/19/2001

CIA-RDP86-00513R002064830004-4"

S/079/62/032/009/011/01). 1048/1242

Synthesis and investigation ...

thosized in the laboratory. Both the rate of condensation and the final yield of butyl acetate increased, while the molecular weight of the polymer decreased with increasing reaction temperature. Thus, at 150°C the reaction was completed within 2 hra, the yield of butyl acetate was 61.4%, and the molecular weight of the polymer was 1290 when the reaction was carried out under atmospheric pressure and 2400 in vacuo. At 180°C the reaction was completed within 1.5 hrs, the yield of butyl acetate was about 75%, and the molecular weight of the polymer was 840 under atmospheric pressure and 1610 in vacuo. The product formed at temperatures above 200°C contained a non-melting, insoluble phase whose structure needs further clarification. The polymer formed at 180°C was a brittle substance, easily soluble in benzene and its homologs, with a softening temperature of 130-140°C. The polymer formed at 150°C was a glass-like, brittle and transparent substance with good adhesion to metals and glass; ita m.p. was 96-100°, and it did not lose its melting characteristics on repeated melting-solidification cycles. There are 2 figures.

SUBMITTED: September 13, 1961 Card 2/2

为新古里面 **在**在在江湖市村,指在北海边被名的东南部。比较多多维亚和克利森和克利森和克斯斯斯亚森,但是自由的维亚斯,但由于一种,由于一种 B/080/62/035/002/006/022 D235/D302 Tolstoguzov, V. B. and Levitskiy, M.M. Gertain properties of oily oligomers of phosphonitrile 15,9209 Zhurnal prikladnoy khimii, v. 35, no. 2, 1962, 290-295 AUTHORS: TEXT: The composition and structure of oligomers of phosphonitrile TITLE S: TEXT: The composition and structure of oligomers of phosphonitric chloride and the properties of high molecular polymeric products of the chained from them are studied who oligomers were obtained by respectively. chloride and the properties of high molecular polymeric products obtained from them are studied. The oligomers were obtained by reacting PCl<sub>5</sub> with NH<sub>4</sub>Cl in a medium of chlorobenzene in the presence PERIODICAL: of quinoline. The trimer and tetramer were removed from the oligomers was mers by extraction with petroleum ether. The yield of oligomers was mers by extraction with petroleum ether within the limits P = 24.96-25.4%, mers by extraction was within the limits P = 24.96-25.4%, mers by extraction was within the limits P = 24.96 and the composition was within the limits P = 24.96 and the composition of molecular weight was 1170 and 12.01-11.74%, Cl. - 59.63.62.5%. The molecular weight 1200 showed a part of the composition of molecular weight was 1170 and titration of a fraction of molecular weight the composition of a fraction of molecular weight was 1170 and titration of a fraction of molecular weight was 1170 and titration of a fraction of molecular weight with petroleum ether. 1) you and titration of a fraction of molecular weight 1200 showed a minimum of three polymer homologues. In order to elucidate the communication the infra-red spectrum in the fraction to infra-red spectrum in the fraction minimum of three polymer nomologues. In order to elucidate the conposition the infra-red spectrum in the frequency range 400 - 3800 Card 1/4 considera si consultanti confide consultanti con consultanti

\$/080/62/035/002/006/022 D235/D302

Certain properties of ...

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cm was studied. The groups P=N, OH and NH were discovered but not P-H or P-OH. The considered oligomers with a degree of polymerization of 10 to 12 have the following structure:

and a portion of the molecule must have the structure

\$/080/62/035/002/006/022 D235/D302

Certain properties of ...

The oligomers were subjected to polymerization in a glass ampoule for 4 hours at 192, 210, 226, 245 and 260°C. On increasing the temperature, consistency of the products changed from a brown paste to a dark brown or black resinous material possessing low adhesion to a dark brown of the samples after extraction with chloroform to glass. Density of the samples after extraction with chloroform was found to be 1.58 - 1.68 at 20°C. The was no relationship between density and temperature of polymerization. A linear relationship was found between the composition of the soluble fraction and the temperature of polymerization which can be written:

 $\frac{M_{sol.fract.}}{M_{sample}}$  x 100 = 175 - 0.57 t°

The authors also studied swelling of the resin and discovered that the degree of swelling decreased with increasing temperature of polymerization. Ageing of the elastomers was considered concluding that the change in weight of the samples during ageing was mainly due to absorption of moisture from the air followed by reaction

Card 3/4

CIA-RDP86-00513R002064830004-4" APPROVED FOR RELEASE: 07/19/2001

Certain properties of ....

S/080/62/035/002/006/022 D235/D302

with the resin with the elimination of HCl; this hydrolysis takes place more quickly the larger the quantity of oily oligomers contained in the resin. The poly-acid formed changes to the more stable tetracompound. There are 6 figures and 5 references: 2 Soviet-bloc and 3 non-Soviet-bloc. The references to the English-language publications read as follows: L. F. Audrieth, R. F. Steedman and A. D. F. Toy, Chem. Rev., 32, 109, 1943; N. L. Paddock, and H. T. Searle, Advances in inorganic chemistry and radiochemistry, 1, 347, 1959; N. L. Paddock, Endeavour., 19, 75, 134, 1960.

SUBMITTED: December 26, 1960

Card 4/4

AUTHOR: Zhivichin, to the control of the control of

TITLE: Synthesis of polymeric propriouse to meet agreets a green

SOURCE: Plasticheskiye massy, no. 7, 1965, 24-26

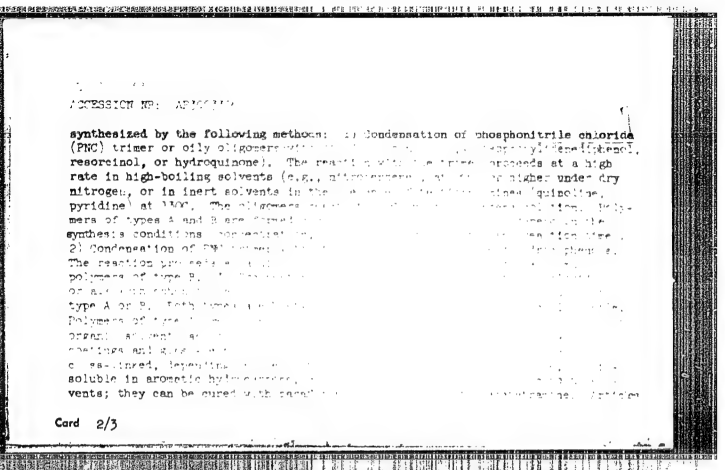
TOPIC TAGS: polyphosphonitrile chloride, alkoxyphosphonitrile chloride, alkoxy-

生性,因为一种,我们是一个人的一个人,我们们是一个人的一个人,这个人的一个人,这个人的一个人的一个人的,我们们的一个人的一个人的人的人的人,我们们的一个人的一个

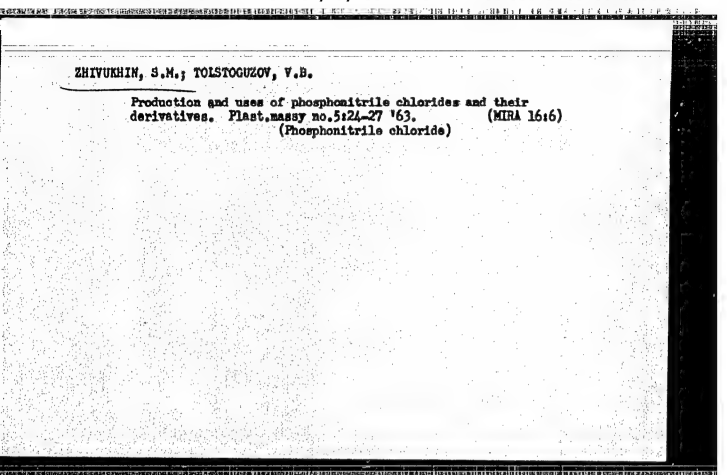
phosphonitrilate, polydroxyarilenephosphonitrilate

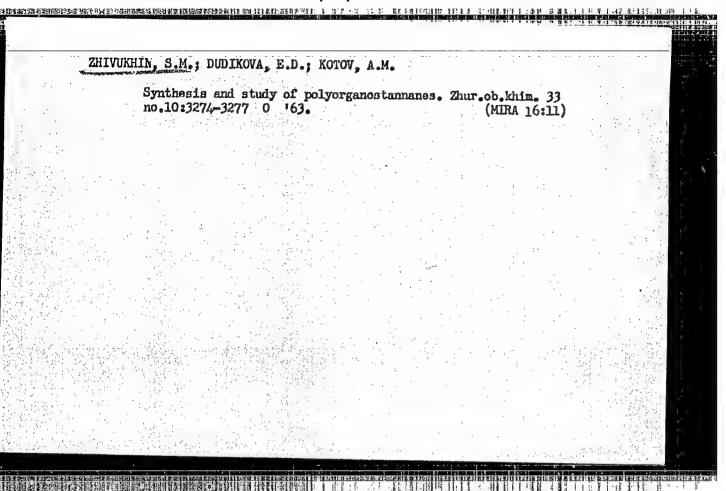
ARSTRACT: Hydrolytically stable polymers of types A and T.

with alternating phosphonitrile and oxygen mention games in the backbone have been Card 1/3



ACCESSION NR: AP3005306 made with polymers of type B exhibit good mechanism! properties but page elseticity and adhesion to metals. The polymer and to use to make molded armoles and glass-reinforced plastics. The presence of incident and incident armoles alkoxy, chloring atoms in phosphonitrile across makes. the adhesion and mechanical properties of the amount of the state of t the and the life to and fire resistance of other polymers can be improved to a training with polymers of type A and B. Orig. art. has: " former ar. ASSOCIATION: none SUBMITTED: 00 DATE ACQ: 30Jul63 ENCL: SUB CODE: CH NO REF SOV: 003 OTHER: 003 Card 3/3 THE PROPERTY OF THE PROPERTY O





ACCESSION NR: AP4041779

\$/0191/64/000/007/0024/0026

AUTHOR: Zhivukhin, S. M., Kireyev, V. V.

TITLE: Some properties of polydihydroxyarylenephosphonitrilates

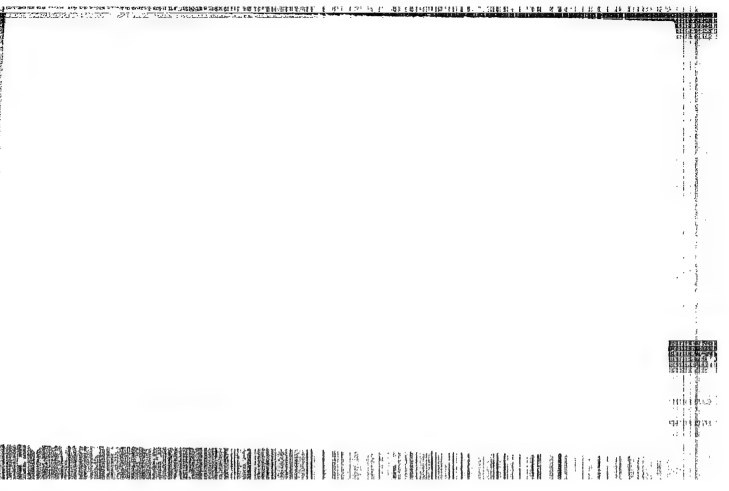
SOURCE: Plasticheskiye massy\*, no. 7, 1964, 24-26

TOPIC TAGS: polydihydroxyarylenephosphonitrilate, phosphonitrile chloride, phenoi diphenylolpropane, resorcinol, hydroquinone, infrared absorption, thermosetting polymer, arylenephosphonitrile, polymer hydrolysis

ABSTRACT: The authors investigated the thermosetting and hydrolytic properties of polydihydroxyarylenephosphonitrilates obtained by the reaction of the trimer phosphonitrile chloride with diatomic phenols, such as diphenylolpropane, resorcingly of and hydroquinone. The infrared absorption spectra are shown and the effects of hardening on the %C, H and Cl in the polymer are tabulated. Thermosetting takes place as a result of the interaction of the functional groups of the polymer and, depending on the temperature, the process can be accompanied by a partial polymerization of the trimer rings. Hydrolysis curves and a study of the products of hydrolysis in either 95% dimethylformamide or 95% acetone at 150 or 55C, respectively, showed that polydihydroxyarylenephosphonitrilates have satisfactory hydrolytic stability. In these polymers, partial or complete substitution of chiorine atoms

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in a fill clic expert	mercal works	vrig. art	• has: 3 1	igures and	2 tables.		
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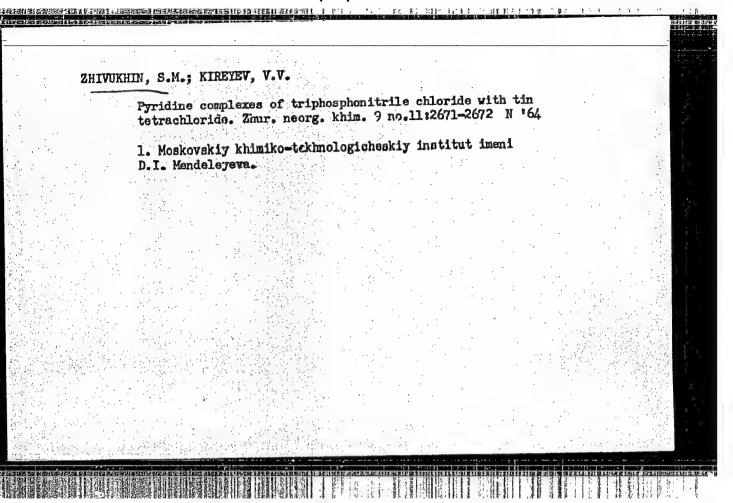


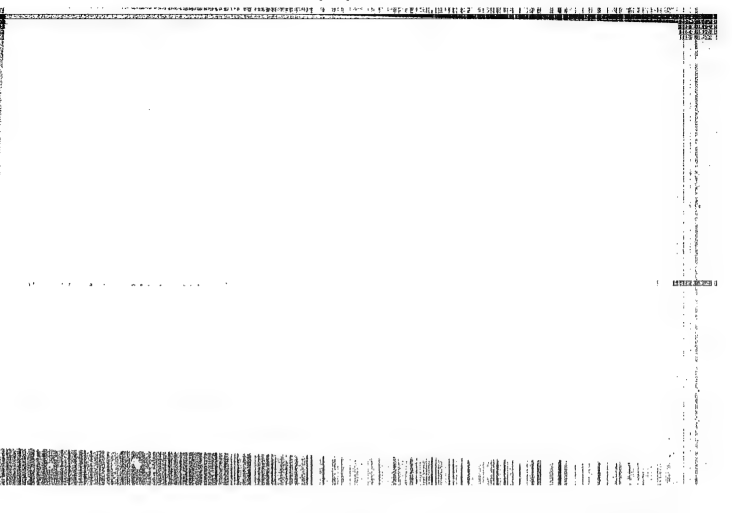


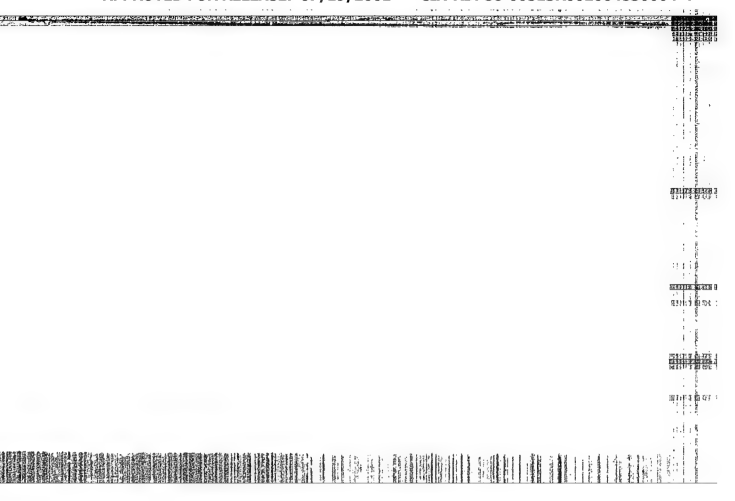


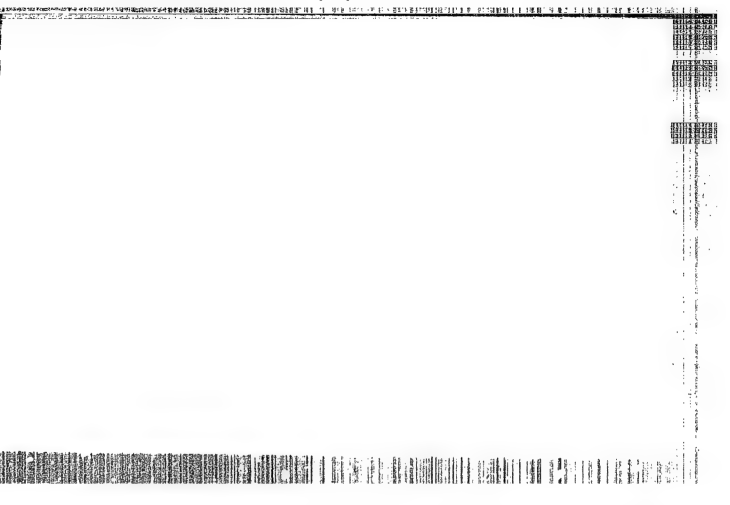
ZHIVUKHIN, S.M.; TOLSTOGUZOV, V.B.; BELYKE, S.I.

Reaction of phosphonitrile chloride with diphenylsilanediol. Zhur.neorg.
(MIRA 17:2)













ZHIVUKHIN, S.M.; TOLSTOGUZOV, V.B.; LUKASHEVSKI, Z.

Reaction of trimeric phosphonitrile chloride with alcoholn and alcoholates. Zhur. neorg. khim, 10 no.7:1653-1656 Jl '65.

(MERA 18:8)

ZHIVUKHIN, S.M.; TOLSTOGUZOV, V.B.; KIREYEV, V.V.; KUZNETHOVA, K.G.

Synthesis of phoaphonitrile chlorides. Zhur. neorg. khim. 10 (MIRA 18:11)

1. Submitted Sopt. 22. 1963.

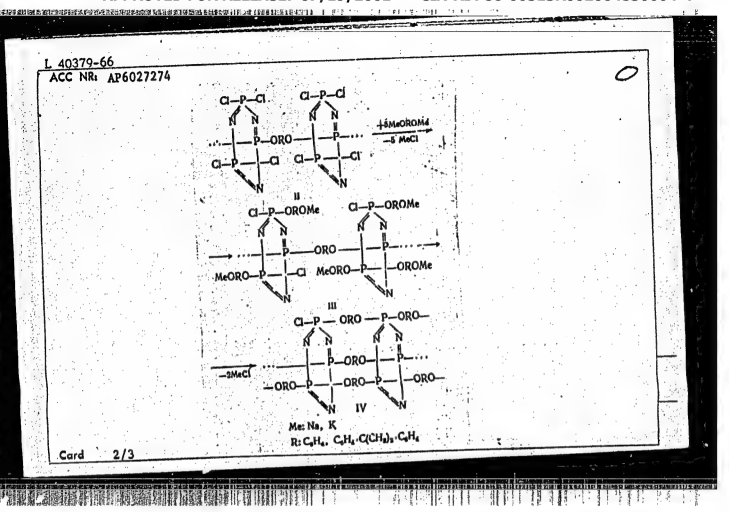
	ACC NR. AP6000974 SQUEETE CODE IN 10286 165 1650 1050 1050
	INVENTOR: Zhivukhin, S. M.; Tolstoguzov, V. B.; Kireyev, V. V. 44,55
	ORG: none
	TITLE: Method for preparing resins. Class 39, No. 17639215
	SOURCE: Byulleten' izobreteniy i tovarnykh znakov, no. 22, 1965, 57
V V	TOPIC TAGS: phosphorus, nesin, polymer, epoxy platic, fire resistant material, thermal stability, lacquer, glass, reinforced plastic, solid machanish property, special-
	ABSTRACT: An Author Certificate has been issued for a preparative method for resins involving phosphorus-containing polymers and epoxy resins/with heating. To obtain polymers with good adhesive and mechanical properties, fire resistance, and thermal stability, polymers consisting of alternating phosphonitrile and hydroxyaromatic structures (polydihydroxyarylenephosphonitrilates) [sic] were used as the phosphorus-containing components. The amount of epoxy resin used does not exceed 50%. The resins obtained are designed for use in lacquer coatings and in glass-reinforced plas-
J.7.1	97/33 [BN]
	SUB CODE: 11,07 / SUBM DATE: 21Jan63 / ATD PRESS: 4/57 UDC: 547.9141
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		ining phosphonitr	ile polymers. C1	ввв 39, No. 1764	23	
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and the statement of the second	the state of the s	in, phosphonitril	programme to the complete management	# "+   10		
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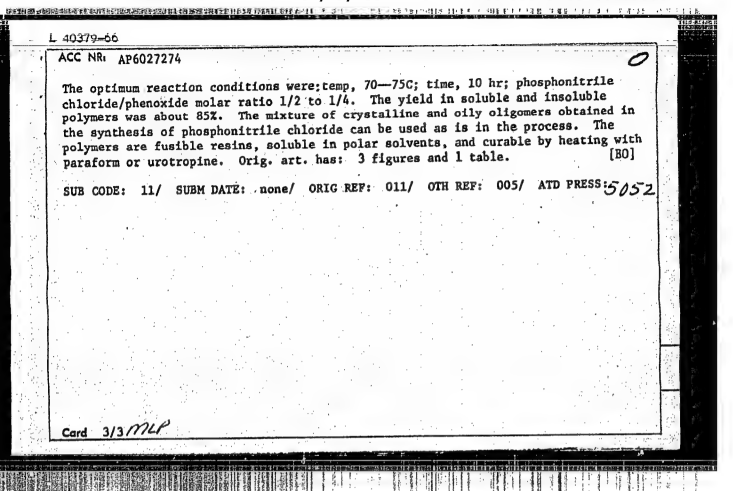
n-F×2	L15324_66EnT(m)/EWP(1)/TYM/RM	
	AUTHORS: Zhivukhin, S. M.; Kireyev, V. V.; Tolstoguzov, V. B.	
	ORG: none	
	TITLE: A method for obtaining phosphonitrile polymers. Class 39, No. 176420	
	SOURCE: Byulleten! izobreteniy i tovarnykh znakov, no. 22, 1965, 61  TOPIC TAGS: polymer, polycondensation, organic phosphorus compound, phosphonitrile	
	ABSTRACT: This Author Certificate presents a method for obtaining phosphonitrile polymers by thermal condensation of phosphonitrile chlorides with dihydroxyphenols. To decrease the condensation temperature, the phenols are used in the form of their alkali metal salts. The condensation is carried out at temperatures not exceeding 150C.	
	SUB CODE: 11/ SUBM DATE: 11Jan63	
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ACC	AP6011238 (A) SOURCE CODE: UR/0413/66/000/006/0076/0076	100
IM	NTOR: Zhivukhin, S. M.; Tolatoguzov, V. B.; Kireyev, V. V.	pr. Y
OR	none	E Court
TI:	E: Preparation of phosphorus-containing polyesters. Class 39,	1
No.	179928	
301 19	CE: Izobreteniya, promyshlennyye obraztsy, tovarnyye znaki, no. 6,	8
TO	C TAGS: polyester, phosphorus containing polymer, transesterification ERIFICATION, PHOSPHORUS, PHOSPHATE, ELYCOL	250
AD	pagm. This Author Certificate introduces a method for preparing	(C) 48
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ph	phorus-containing polyesters by transesterification of phosphates	
ph wi al	phorus-containing polyesters by transesterification of phosphates of glycols. To extend the variety of <u>fire-resistant</u> wolfied additives oxyphosphonitrilates and/or <u>alkoxyphosphonitrile</u> chlorides are	5.0
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pho wi all su SU	phorus-containing polyesters by transesterification of phosphates a glycols. To extend the variety of <u>fire-resistant</u> iodified additives exphosphonitrilates and/or <u>alkoxyphosphonitrile chlorides</u> are [LD]	at to the traffic administration of

ACC NR: AP6027274 (A) SOURCE CODE: UR/0191/66/000/008/0015/0018  AUTHOR: Zhivukhin, S. M.; Kireyev, V. V.; Tikhonova, G. S. 33  ORG: none  TITLE: Polymers pased on phosphonitrile chlorides and bisphenol phenoxides  SOURCE: Plasticheskiye massy, no. 8, 1966, 15-18  TOPIC TAGS: phosphorus containing nalymar, phosphonitrile chloride, bisphenol phenoxide, best rosistant polymer, reactive polymer, Polymer HEAT RESISTANCE, PHENOL, RESIN, POLYMER CHEMISTRY  ABSTRACT: A study has been made of the synthesis of phosphonitrile chloride-bisphenol phenoxide polymers. This reaction is of interest because it has the following advantages: it proceeds rapidly at 70-130C in the absence of catalysts without the liberation of HCL. The products exhibit high heat resistance and contain reactive phenoxide end groups. The starting materials were phosphonitrile chloride trimer and/or phosphonitrile chloride oily oligomers, and Na or K phenoxides of 2,2-bis(p-hydroxylphenyl) propane or of resorcinal. The reactions were conducted in dehydrated m-xylene, dioxane, or methyl ethyl ketone. The procedure is described in the source. The following reaction mechanism is proposed



APPROVED FOR RELEASE: 07/19/2001 CIA-RDP86-00513R002064830004-4"



ACC NR	AP6012719 (A) SOURCE CODE: UR/0190/66/008/004/0727/0731
AUTHOR:	: Zhivukhin, S. M.; Tolstoguzov, V. B.; Yakobson, F. I.
ORG: M	Moscow Institute of Chemical Technology im. D. I. Mendeleyev (Moskovskiy khimiko- logicheskiy institut)
TITLE:	Synthesis of polydioxyarylenephosphonitrilates 1
SOURCE:	Vysokomolekulyarnyye soyedineniya, v. 8, no. 4, 1966, 727-731
TOPIC T	PAGS: phosphonitrilate, esterification, polyesterification
diatomi The rat and on ester lower m to 10,0 aromati authors	T: Polyester exchange reaction of hexabutoxytriphosphonitrilate with c phenols was carried out at the molar component ratio from 1:1 to 1:1.5. See of polyester exchange depends on the concentration of the reaction mixture the type of diatomic phenol. Resorsinal is somewhat more active in polyexchange. In the case of hydroquinone, products with higher substitution but colecular weight are formed. Products which have a molecular weight of 3000 loo are easily soluble in alcohols and ketones and partially soluble in a c and aliphatic solvents. Orig. art. has: 5 figures and 2 tables. [Based on [NT]]
SUB COD	DE: 11, 07/ SUBM DATE: 09May65/ ORIG REF: 003/ OTH REF: 003
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* : !=+		L 15772-66 EVT(m)/EVP(1)/T/ETC(m)-6 - WW/RM SOURCE CODE: UR/0080/66/039/001/0234/0237	
		AUTHOR: Zhivukhin, S. M.; Kireyev, V. V.; Zelenetskiy, A. N. B  ORG: none	
		TITLE: The reaction of phosphonitrile chloride trimor with dihydric phenols  SOURCE: Zhurnal prikladnoy khimii, v. 39, no. 1, 1966, 234-237	
		ABSTRACT: Polymers derived from phosphonitrile chlorides and dihydric phenols have high thermal stability fire resistance and other desirable properties. The purpose of this work was to investigate the reaction between phosphonitrile chloride trimer and 2,2-bis-(p-hydroxyphenyl)propane, resortinol and hydroquinone. It was found that phosphonitrile chloride trimer does not react with dihydric phenols below 180C, lither in the melt or in organic solvents. The reaction, accompanied by liberation of HCl, takes place at 200C, or above, in nitro-	
1 1 1 1 1 1 1		benzene or ditolylmethane, or at lower temperatures in some organic solvents in the presence of quinoline or pyridine. The effect of the duration of the reaction, and of the ratio and concentration of starting materials on the composition of Card 1/2  UDC: 547.56+541.6	
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L 15772-66	
ACC NR: AP6005522  the product is shown. Some conclusions concerning the structure of	the product
are drawn from elemental and functional-group analyses, molecular we infrared spectra. Orig. art. has: 4 figures and I table.  SUB CODE: 11,07 SUBM DATE: Olju163/ ORIG REF: 001/ OTH REF:	(vs)
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ZHIVUKHINA, G. M. Cand Biol Soi -- "Frost resistance and state of rest of winter wheat under conditions of Moskovskaya Oblast." Mos, 1981 (Mosy Oblast Ped Inst im N. K. Krupskaya). (KL, 4-61, 192)

-125-

17(1)
AUTHORS: Genkel', P. A., Zhivukhina, G. M. SOV/20-127-1-61/65

TITLE: The Process of Protoplasm Isolation as the Second Phase of Winter Wheat Hardening (Protsess obosobleniya protoplazmy kak vtoraya faza zakalivaniya ozimykh pshenits)

PERIODICAL: Doklady Akademii nauk SSSR, 1959, Vol 127, Nr 1, pp 220-223 (USSR)

The plants resistant to frost are characterized by a period of long-lasting and profound rest. The resistance of the tissues of resting plants is determined on the whole by the conditional peculiarities of the plasma (Refs 1-3). The lacking of growth processes (Refs 4-8 for winter wheat), the reduced metabolism intensity as well as the protoplasm separation are characteristic of the period of rest. During the period of rest the content in growth substances is considerably reduced (Refs 9-12). In plants hardened against frost and in a state of rest the protoplasm is characterized by a higher viscosity and by a reduced permeability (Refs 18-19). The physiological processes are of little intensity in winter wheat (Refs 20, 21). The state of rest is of little stability in winter crops. However, if winter wheat is stored during the winter in a warm room for a few days only.

Card 1/4

ABSTRACT:

The Process of Protoplasm Isolation as the Second Phase of Winter Wheat Hardening

SOV/20-127-1-61/65

the growth processes start quickly in contrast to ligneous plants. The period of rest of the winter crops is very short because no profound transformation of the protective-and supply substances takes place in their cells and because they have no reserve fat which guarantees a stable state of rest (Ref 3). The transition to the state of rest takes place under the influence of autumn conditions of temperature and light. Under the same circumstances the hardening of the plants against low temperature takes place. Thus the transition to the period of rest and the hardening occur during the same period in the life of plants, and both reflect the same processes during the course of which the plants attain the resistance to frost. The hardening is attained in two stages: 1) accumulation of carbohydrates; 2) change of the physico-chemical protoplesm properties (Ref 2). In the present paper the cyto-physiological state of the winter wheat was investigated. The observations of the authors confirmed that the protoplasm becomes peeled off from the cell-coverings in the late autumn- and winter period. The degree of the state of rest differs in the individual organs of the winter wheat. The state of rest of the leaves

Card 2/4

The Process of Protoplasm Isolation as the Second Phase of Winter Wheat Hardening

SOV/20-127-1-61/65

is not long. The protoplasm is partly detached only in a few epidermic cells. In autumn and winter, a concave plasmolysis is predominant. At the turn of the year many damaged and dead cells appear. The most complete rest (according to the number (according to the length of separation) were found in the growth cone and in the tillering knots (Fig 1). The observations the authors showed: 1) that the protoplasm separation from that is the accumulation of the soluble carbohydrates in the In other words, the conditions of the separation process of second stage of hardening of the winter crops is attained which is connected with the physico-chemical changes of the

Card 3/4

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The Process of Protoplasm Isolation as the Second Phase of Winter Wheat Hardening

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protoplasm. The said separation occurs obviously just in the second hardening stage and forms the final stage of the development of the resistance to frost of the winter crops under autumn conditions. There are 1 figure, 1 table, and 22 references, 21 of which are Soviet.

ASSOCIATION:

Institut fiziologii rasteniy im. K. A. Timiryazeva Akademii nauk SSSR (Institute of Plant Physiology imeni K. A. Timiryazev of the Academy of Sciences. USSR)

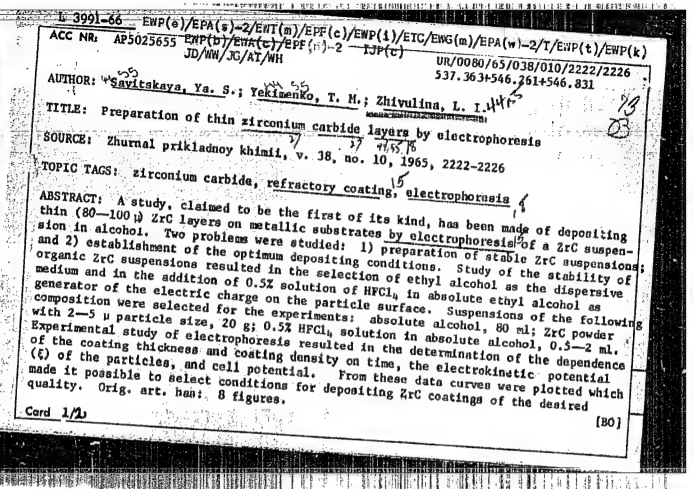
PRESENTED:

February 2, 1959, by A. L. Kursanov, Academician

SUBMITTED:

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